AMENDMENTS TO THE CLAIMS

1. (Currently amended) A coated implant for in vivo-anchoring to a biological tissue or another implant, which coated implant comprises an implant having a pre-treated surface and on said pre-treated surface one or more layers of mainly non-hydrated chemically bonded ceramic material, characterised in that

each layer of said ceramic material independently comprises a first binder phase selected from the group consisting of aluminates, silicates, phosphates, sulphates and combinations thereof, and that said ceramic material is chemically and/or mechanically bound to said implant, and in that

the coated implant additionally is coated with a ceramic paste comprising a powdered calcium-based binder of aluminate and/or silica and a hydration liquid.

- 2. (Original) A coated implant according to claim 1, characterised in that the first binder phase comprises cations selected from the group consisting of Ca, Sr and Ba.
- 3. (Original) A coated implant according to claim 2, characterised in that the cations are Cacations.
- 4. (Original) A coated implant according to claim 3, characterized in that the first binder phase comprises calcium aluminates.
- 5. (Original) A coated implant according to claim 4, characterized in that the first binder phase comprises one or more of the phases 3CaO·Al₂O₃, 12CaO·7Al₂O₃ CaO·Al₂O₃, CaO·Al₂O₃ and CaO·6Al₂O₃.
- 6. (Currently amended) A coated implant according to claim 1, characterised in that the ceramic material further comprises water-soluble phosphate or a phase (such as a phophate salt) that has the capacity to form water-soluble phosphate.

- 7. (Previously presented) A coated implant according to claim 1, characterised in that said one or more non-hydrated layers have a porosity below 50%.
- 8. (Previously presented) A coated implant according to claim 1, characterised in that the surface roughness of the pre-treated surface of the implant has a Ra-value of less than 10 μ m, but not smaller than 0.5 μ m.
- 9. (Currently amended) A coated implant according to claim 1, characterised in that the number of layers of the coating is 1-5, in addition to the paste layer.
- 10. (Previously presented) A coated implant according to claim 1, characterised in that an innermost layer has a thickness in the interval from nanometer level to less than 10 μm.
- 11. (Currently amended) A coated implant according to claim 1, characterised in that an outermost layer beneath the paste has a surface treated to a surface roughness of Ra<20 μ m, but not smaller than 0.5 μ m.
- 12. (Previously presented) A coated implant according to claim 1, characterised in that it comprises at least two layers and that each layer outside the innermost one independently has a thickness of less than 50 μ m, but not smaller than 5 μ m.
- 13. (Currently amended) A coated implant according to claim 1, characterised in that said implant is a medical, orthopaedic or dental implant, such as an artificial orthopaedic device, a spinal implant, a joint implant, an attachment element, a bone nail, a bone screw, and a bone reinforcement plate.
- 14. (Previously presented) A coated implant according to claim 1, characterised in that said implant is of a ceramic, metallic or polymeric material.

- 15. (Original) A coated implant according to claim 14, characterised in that said implant material has been selected from titanium, stainless steels, alumina, zirconia and medical grade plastics.
- 16. (Previously presented) A coated implant according to claim 1, characterised in that the implant surface is oxidized.
- 17. (Original) A coated implant according to claim 16, characterised in that said oxide is a double oxide of titanate, silicate or aluminate type.
- 18. (Previously presented) A coated implant according to claim 1, characterised in that said mechanical binding to the implant is achieved by sub-micron size crystallites of hydrates precipitated on the surface of said implant.
- 19. (Original) A coated implant according to claim 18, characterised in that the crystallite size is less than 100 μm .
- 20. (Previously presented) A coated implant according to claim 1, characterised in that the powdered mainly non-hydrated ceramic material has a particle size of 0.1 to 20 μm.
- 21. (Withdrawn) A method of manufacturing a coated implant according to claim 1, which method comprises the steps of:
 - -pre-treating the surface of an implant,
- -applying on said pre-treated surface one or more layers of mainly powdered non-hydrated ceramic material, which layers independently comprises a first binder phase selected from the group consisting of aluminates, silicates, phosphates, sulphates and combinations thereof, and
- -optionally pre-hydrating said ceramic material by contacting it with a curing liquid or body fluid,
- -thereby forming a chemical and/or mechanical bond between the ceramic material and said implant.

- 22. (Withdrawn) A method according to claim 21, characterised in that said pre-treatment is selected from a group consisting of oxidation including low-temperature oxidation, thermal treatment including solid state diffusion and ion bombarding, etching including the use of salt melts, calcination, sand-blasting and grinding.
- 23. (Withdrawn) A method according to claim 21, characterised in that the surface roughness of the implant after pre-treatment has a Ra-value of less than 10 µm, but not smaller than 0.5 µm.
- 24. (Withdrawn) A method according to claim 23, characterised in that the innermost layer of the coating is applied on the implant surface by any of the following techniques: thermal spraying, flame spraying, Electro Deposition Spraying (EDS), plasma spraying, dipping and spin coating.
- 25. (Withdrawn) A method according to claim 23, characterised in that when the surface roughness of the implant has a Ra-value of less than 1μm, but not smaller than 0.05 μm, the innermost layer of the coating is applied on the implant surface by any of the following techniques: Chemical Vapor Deposition (CVD), Physical Vapor Deposition (PVD), laser techniques including laser cladding, Electrolytic Deposition (ED), and sol-gel techniques.
- 26. (Withdrawn) A method according to any of claims 25, characterised in that when the coating only comprises one layer, said layer is applied using Physical Vapor Deposition (PVD).
- 27. (Withdrawn) A method according to claim 21, characterised in that said one or more layers of the coating are thinned, preferably by a process selected from the group consisting of grinding, sand blasting, dry etching and chemical treatment including dissolution.
- 28. (Withdrawn) A method according to claim 27, characterised in that in connection with said thinning, a partial densification of said one or more layers is performed, preferably by drying up of particles and precipitation including sol-gel techniques.

- 29. (Withdrawn) A method according to claim 21, characterised in that the pre-hydration is performed by dipping, spraying, spin coating or tape casting the coated implant in/with such an additional hydration liquid.
- 30. (Withdrawn) A method according to claim 21, characterised in that the powdered, mainly non-hydrated ceramic material, has a particle size of 0.1 to 20 µm.
- 31. (Withdrawn) A ceramic paste, characterised in that it comprises a powdered calcium-based binder of aluminate and/or silicate and a hydration liquid.
- 32. (Withdrawn) A ceramic paste according to claim 31, characterised in that it has the form of granules of a size below 1 mm and a granule compaction density above 35 %.
- 33. (Withdrawn) A ceramic paste according to claim 32, characterised in that the granules have a mean size of at least 30 μ m, but 250 μ m at the most.
- 34. (Withdrawn) A ceramic paste according to claim 31, characterised in that it comprises an organic additive, preferably a hydrophilic polyacrylic and/or polycarboxylate compound.
- 35. (Withdrawn) An implantation kit for in vivo-anchoring an implant to a biological tissue or another implant, comprising the coated implant according to claim 1 and optionally a curing liquid capable of hydrating the binder phase of the coated implant and a paste according to claim 31, wherein the ceramic powder and hydration liquid of the paste are kept separately.
- 36. (New) A coated implant according to claim 6, wherein the ceramic material further comprises a phosphate salt.
- 37. (New) A coated implant according to claim 13, wherein said implant is an artificial orthopaedic device, a spinal implant, a joint implant, an attachment element, a bone nail, a bone screw, or a bone reinforcement plate.